## Hepatoprotective effects of systemic ER activation ChIPseq/Epigenome genome - Annotation of Peak files and feature examination

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Load the enhancers and promoters which were demarcated using various histone modifications. Then annotate these features and export them as tables. For promoters, only one peak per gene, the closest one, is permitted, multiple TSS are disregaded.

```
library("ChIPpeakAnno")
library("GenomicRanges")
options(connectionObserver = NULL) #That is a work-around, as the org.Mm. package cannot be loaded
library("org.Mm.eg.db")
library("biomaRt")
library("tidyverse")
import_list <- list(</pre>
promoters_genomewide <- read.delim("results/Epigenome_analysis/Final_promoter_files/prom.3.genomewide.F
enhancers_genomewide <- read.delim("results/Epigenome_analysis/Final_enhancer_files/enh.5.FINAL.genomew
enhancers_allDB <- read.delim("results/Epigenome_analysis/Final_enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enh.5.FINAL.H3K27ac_broadens.enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_files/enhancer_fil
promoters_allDB <- read.delim("results/Epigenome_analysis/Final_promoter_files/prom.8.FINAL.H3K27ac_bro
enhancers_HFDup <- read.delim("results/Epigenome_analysis/Final_enhancer_files/enh.7.HFDup.FINAL.H3K27a
promoters_HFDup <- read.delim("results/Epigenome_analysis/Final_promoter_files/prom.16.FINAL.HFDup.H3K2</pre>
enhancers_HFDdown <- read.delim("results/Epigenome_analysis/Final_enhancer_files/enh.9.HFDdown.FINAL.H3
promoters HFDdown <- read.delim("results/Epigenome analysis/Final promoter files/prom.24.FINAL.HFDdown."
names(import_list) <- c("promoters_genomewide", "enhancers_genomewide", "enhancers_allDB", "promoters_a
##Filter out all locations which are NOT on a chromosome.
import_list2 <- list()</pre>
for (i in names(import_list)) {
object <- import_list[[i]] # Save the respective dfs in a non-list object, and put that one into a list
object <- dplyr::filter(object, grepl("chr", V1))</pre>
import_list2[[i]] <- object</pre>
#listEnsemblArchives()
mart <- useMart(biomart = "ensembl", dataset = "mmusculus_gene_ensembl", host = "https://sep2019.archiv</pre>
# use the qetAnnotation function to obtain the TSS for ensembl GRCh38.p13.
annoData <- getAnnotation(mart, featureType = "TSS")</pre>
```

```
# Annotate the peak files.
peaks_annotated <- list()</pre>
for (i in names(import_list2)) {
object <- import_list2[[i]]</pre>
colnames(object) <- c("chrom", "start", "end")</pre>
nrow(object)
object2 <- makeGRangesFromDataFrame(object, start.field = "start", end.field = "end", ignore.strand = '
#Give ranges numeric names in order
names(object2) <- c(1:length(object2))</pre>
#Annotate granges with the nearest TSS
object3 <- annotatePeakInBatch(object2,</pre>
                                AnnotationData=annoData,
                                featureType = "TSS",
                                output="nearestLocation",
                                PeakLocForDistance = "start")
object3 <- as.data.frame(object3)</pre>
peaks_annotated[[i]] <- object3</pre>
# For the promoters, remove the duplicated gene names and only allow the closest peak to a given gene.
library(data.table)
peaks_annotated[["promoters_genomewide"]] <- peaks_annotated[["promoters_genomewide"]] %>% dplyr::group
peaks_annotated[["promoters_genomewide"]] <- setDT(peaks_annotated[["promoters_genomewide"]])[order(sho</pre>
peaks_annotated[["promoters_HFDdown"]] <- peaks_annotated[["promoters_HFDdown"]] %>% dplyr::group_by(fe
peaks_annotated[["promoters_HFDdown"]] <- setDT(peaks_annotated[["promoters_HFDdown"]])[order(shortestD</pre>
peaks_annotated[["promoters_HFDup"]] <- peaks_annotated[["promoters_HFDup"]] %>% dplyr::group_by(featur
peaks_annotated[["promoters_HFDup"]] <- setDT(peaks_annotated[["promoters_HFDup"]])[order(shortestDista</pre>
peaks_annotated[["promoters_allDB"]] <- peaks_annotated[["promoters_allDB"]] %>% dplyr::group_by(featur
peaks annotated[["promoters allDB"]] <- setDT(peaks annotated[["promoters allDB"]])[order(shortestDista</pre>
saveRDS(peaks_annotated, "results/Epigenome_analysis/annotated_diffbind_and_genomewide_promoters_enhanc
```

## Export the bedfiles, which are required for further analyses

```
peaks_annotated_bed <- list()
for (i in names(peaks_annotated)) {
   peaks_annotated_bed[[i]] <- peaks_annotated[[i]] %>% dplyr::select("chrom"=seqnames, start, end)
write.table(peaks_annotated_bed[[i]], paste0("results/Epigenome_analysis/",i, "_", as.character(length())
```

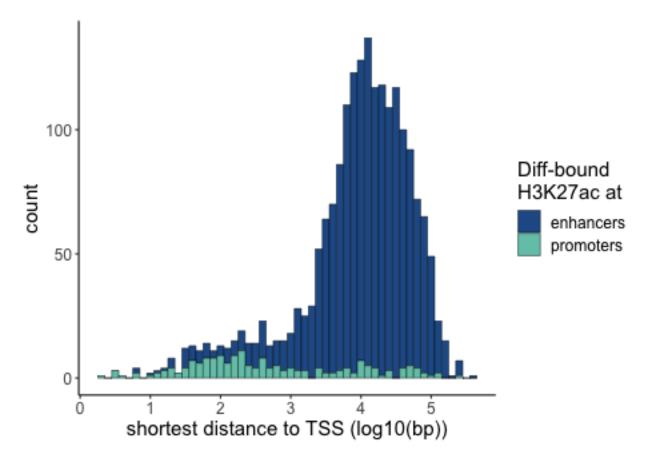
}

Summary statistics about distance to nearest TSS to evaluate how well the enhancers/promoters were determined.

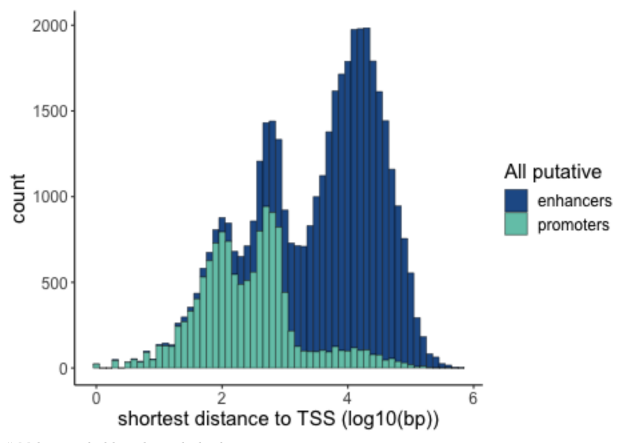
```
# Print the distance metrics
distance_metics <- list()</pre>
for (i in names(peaks_annotated)) {
  print(paste("The median / mean / min / max for", i, "are:"))
print(
summary(peaks_annotated[[i]]$shortestDistance))
  distance_metics[[i]] <- summary(peaks_annotated[[i]]$shortestDistance)</pre>
}
## [1] "The median / mean / min / max for promoters_genomewide are:"
      Min. 1st Qu. Median
##
                              Mean 3rd Qu.
                                              Max.
##
                74
                       245
                              2451
                                       670 395073
## [1] "The median / mean / min / max for enhancers_genomewide are:"
##
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                             22274
                     11200
##
         0
              3533
                                     27152 617593
  [1] "The median / mean / min / max for enhancers_allDB are:"
##
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
                     12894
                             25051
##
              5242
                                     32282 437429
## [1] "The median / mean / min / max for promoters_allDB are:"
##
       Min. 1st Qu.
                       Median
                                  Mean 3rd Qu.
                76.0
##
                        253.5 10036.8
                                        5304.0 256716.0
## [1] "The median / mean / min / max for enhancers HFDup are:"
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                               Max.
##
##
         5
              4949
                     11838
                             23426
                                     29252 281639
  [1] "The median / mean / min / max for promoters HFDup are:"
##
               1st Qu.
                                      Mean
                                             3rd Qu.
##
       Min.
                          Median
##
        1.00
                 86.25
                          237.50 10915.61
                                             4595.50 256716.00
## [1] "The median / mean / min / max for enhancers_HFDdown are:"
      Min. 1st Qu. Median
                              Mean 3rd Qu.
##
              6119
                             28302
##
                     15100
                                     37891 437429
## [1] "The median / mean / min / max for promoters_HFDdown are:"
            1st Qu.
##
                       Median
                                  Mean 3rd Qu.
       Min.
##
                62.5
                        307.0
                                         5304.0 126490.0
                                8865.1
# Add a column to identify the regions
promoters_anno <- peaks_annotated$promoters_allDB %>% mutate(element = "promoters")
enhancers_anno <- peaks_annotated$enhancers_allDB %>% mutate(element = "enhancers")
promoters_genomewide_anno <- peaks_annotated$promoters_genomewide %>% mutate(element = "promoters")
enhancers_genomewide_anno <- peaks_annotated$enhancers_genomewide %>% mutate(element = "enhancers")
# Row-bind the dataframes together
combined DBsites anno <- rbind(promoters anno, enhancers anno)
combined_allSites_anno <- rbind(promoters_genomewide_anno, enhancers_genomewide_anno)
```

```
combined_DBsites_anno$shortestDistance <- combined_DBsites_anno$shortestDistance +1
combined_allSites_anno$shortestDistance <- combined_allSites_anno$shortestDistance +1

# Plot the histograms
ggplot(combined_DBsites_anno)+
   geom_histogram(aes(x=log10(shortestDistance), fill=factor(element)), binwidth=0.1, color="black", siz theme_classic() +
   scale_fill_manual("Diff-bound\nH3K27ac at", values = c("#235b95", "#73c6b6")) +
   theme(text=element_text(size=15)) +
   xlab("shortest distance to TSS (log10(bp))")</pre>
```



```
ggplot(combined_allSites_anno)+
  geom_histogram(aes(x=log10(shortestDistance), fill=factor(element)), binwidth=0.1, color="black", siz
  theme_classic() +
  scale_fill_manual("All putative", values = c("#235b95", "#73c6b6")) +
  theme(text=element_text(size=15)) +
  xlab("shortest distance to TSS (log10(bp))")
```

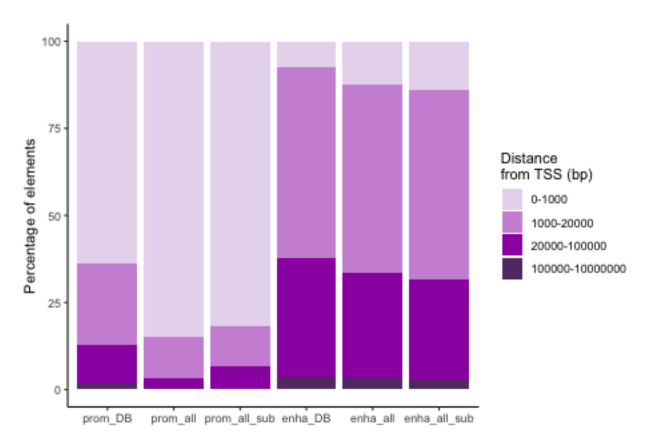


# Make a stacked bar plot with the distances.

```
# set up cut-off values
breaks \leftarrow c(0,1000,20000,100000,10000000)
# specify interval/bin labels
tags <- c("[0-1000)","[1000-20000)", "[20000-100000)", "[100000-10000000)")
# bucketing values into bins
promoters_anno_bins <- cut(peaks_annotated$promoters_allDB$shortestDistance,</pre>
                  breaks=breaks,
                  include.lowest=TRUE,
                  right=FALSE,
                  labels=tags)
enhancers_anno_bins <- cut(peaks_annotated$enhancers_allDB$shortestDistance,</pre>
                  breaks=breaks,
                  include.lowest=TRUE,
                  right=FALSE,
                  labels=tags)
promoters_genomewide_anno_bins <- cut(peaks_annotated$promoters_genomewide$shortestDistance,
                  breaks=breaks,
                  include.lowest=TRUE,
                  right=FALSE,
                  labels=tags)
enhancers_genomewide_anno_bins <- cut(peaks_annotated$enhancers_genomewide$shortestDistance,
                  breaks=breaks,
```

```
include.lowest=TRUE,
                  right=FALSE,
                  labels=tags)
set.seed(500)
promoters_genomewide_anno_random_182 <- promoters_genomewide_anno[sample(nrow(promoters_genomewide_anno
promoters_genomewide_anno_random_182_bins <- cut(promoters_genomewide_anno_random_182$shortestDistance,
                  breaks=breaks,
                  include.lowest=TRUE,
                  right=FALSE,
                  labels=tags)
enhancers_genomewide_anno_random_1816 <- enhancers_genomewide_anno[sample(nrow(enhancers_genomewide_anno
promoters_genomewide_anno_random_1816_bins <- cut(enhancers_genomewide_anno_random_1816$shortestDistanc
                  breaks=breaks,
                  include.lowest=TRUE,
                  right=FALSE,
                  labels=tags)
# inspect bins. The occurrence of each bin is counted.
summary_distance <- data.frame(prom_DB = summary(promoters_anno_bins),</pre>
           enha_DB = summary(enhancers_anno_bins),
           prom_all = summary(promoters_genomewide_anno_bins),
           prom_all_sub = summary(promoters_genomewide_anno_random_182_bins),
           enha_all = summary(enhancers_genomewide_anno_bins),
           enha_all_sub = summary(promoters_genomewide_anno_random_1816_bins))
# Getting the colsums for number of elements
summary_distance.mat <- as.matrix(summary_distance)</pre>
colsums <- colSums(summary_distance[1:6])</pre>
# .. and normalize to the colSums to get percentages
summary_distance.mat <- as.data.frame(round((</pre>
  sweep(summary_distance.mat,2,colsums, "/")*100)
  ,<mark>2</mark>))
summary_distance <- tibble::rownames_to_column(summary_distance.mat)</pre>
# long format needed for plotting
summary_distance_long <- pivot_longer(summary_distance, cols = 2:7) %>%
  group_by(name)
# Replacing the brackets
summary_distance_long$rowname <- gsub(pattern="\\[", "", summary_distance_long$rowname)</pre>
summary_distance_long$rowname <- gsub(pattern="\\)", "", summary_distance_long$rowname)</pre>
# Setting the order for plotting
order_dist <- c("0-1000", "1000-20000", "20000-100000", "100000-10000000")
order_element <- c("prom_DB", "prom_all", "prom_all_sub", "enha_DB", "enha_all", "enha_all_sub")</pre>
# Plotting the stacked bar plots
ggplot(summary_distance_long) +
  geom_col(aes(y=value, x=factor(name, levels=order_element), fill=factor(rowname, levels=order_dist)))
```

```
scale_fill_manual("Distance\nfrom TSS (bp)", values=c("#e8daef","#ce93d8", "#9c27b0","#633974")) +
ylab("Percentage of elements") +
xlab("") +
theme(text=element_text(size=15)) +
theme_classic()
```



## sessionInfo()

```
## R version 4.0.3 (2020-10-10)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Big Sur 10.16
## Matrix products: default
         /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
## [1] stats4
                                    graphics grDevices utils
                parallel stats
                                                                  datasets
## [8] methods
##
## other attached packages:
  [1] data.table_1.14.0
                            forcats_0.5.1
                                           stringr_1.4.0
```

```
## [4] dplyr_1.0.6
                             purrr_0.3.4
                                                  readr_1.4.0
## [7] tidyr_1.1.3
                             tibble_3.1.2
                                                  ggplot2_3.3.3
## [10] tidyverse_1.3.1
                             biomaRt 2.46.3
                                                  org.Mm.eg.db_3.12.0
## [13] AnnotationDbi_1.52.0 Biobase_2.50.0
                                                  ChIPpeakAnno_3.24.2
## [16] GenomicRanges_1.42.0 GenomeInfoDb_1.26.7
                                                  IRanges_2.24.1
## [19] S4Vectors 0.28.1
                             BiocGenerics 0.36.1
## loaded via a namespace (and not attached):
##
     [1] colorspace_2.0-1
                                     ellipsis_0.3.2
##
     [3] futile.logger_1.4.3
                                     XVector_0.30.0
##
     [5] fs_1.5.0
                                     rstudioapi_0.13
##
     [7] farver_2.1.0
                                     bit64_4.0.5
##
     [9] fansi_0.5.0
                                     lubridate_1.7.10
##
  [11] xml2_1.3.2
                                     splines_4.0.3
##
  [13] cachem_1.0.5
                                     knitr_1.33
##
    [15] jsonlite_1.7.2
                                     Rsamtools_2.6.0
##
  [17] broom_0.7.6
                                     dbplyr_2.1.1
                                     graph_1.68.0
##
  [19] png_0.1-7
##
  [21] compiler_4.0.3
                                     httr_1.4.2
   [23] backports 1.2.1
                                     assertthat 0.2.1
## [25] Matrix_1.3-3
                                     fastmap_1.1.0
                                     cli_3.2.0
## [27] lazyeval_0.2.2
## [29] formatR 1.9
                                     htmltools 0.5.1.1
## [31] prettyunits_1.1.1
                                     tools 4.0.3
## [33] gtable_0.3.0
                                     glue_1.6.2
## [35] GenomeInfoDbData_1.2.4
                                     rappdirs_0.3.3
## [37] Rcpp_1.0.6
                                     cellranger_1.1.0
## [39] vctrs_0.3.8
                                     Biostrings_2.58.0
## [41] multtest_2.46.0
                                     rtracklayer_1.50.0
## [43] xfun_0.31
                                     rvest_1.0.0
## [45] lifecycle_1.0.0
                                     ensembldb_2.14.0
##
  [47] XML_3.99-0.6
                                     zlibbioc_1.36.0
##
  [49] MASS_7.3-53.1
                                     scales_1.1.1
## [51] BSgenome_1.58.0
                                     hms_1.1.0
   [53] MatrixGenerics 1.2.1
                                     ProtGenerics_1.22.0
## [55] SummarizedExperiment_1.20.0 RBGL_1.66.0
## [57] AnnotationFilter 1.14.0
                                     lambda.r 1.2.4
## [59] yaml_2.2.1
                                     curl_4.3.1
##
   [61] memoise_2.0.0
                                     stringi_1.6.2
## [63] RSQLite_2.2.6
                                     highr_0.9
## [65] GenomicFeatures 1.42.3
                                     BiocParallel_1.24.1
                                     pkgconfig_2.0.3
## [67] rlang_0.4.11
## [69] matrixStats_0.58.0
                                     bitops_1.0-7
## [71] evaluate_0.14
                                     lattice_0.20-41
## [73] labeling_0.4.2
                                     GenomicAlignments_1.26.0
## [75] bit_4.0.4
                                     tidyselect_1.1.1
## [77] magrittr_2.0.1
                                     R6_2.5.0
##
  [79] generics_0.1.0
                                     DelayedArray_0.16.3
## [81] DBI_1.1.1
                                     withr_2.4.2
## [83] pillar_1.6.1
                                     haven_2.4.1
## [85] survival_3.2-10
                                     KEGGREST_1.30.1
## [87] RCurl_1.98-1.3
                                     modelr 0.1.8
## [89] crayon_1.4.1
                                     futile.options_1.0.1
## [91] utf8_1.2.1
                                     BiocFileCache 1.14.0
```

```
## [93] rmarkdown_2.14 progress_1.2.2
## [95] grid_4.0.3 readxl_1.3.1
## [97] blob_1.2.1 reprex_2.0.0
## [99] digest_0.6.27 VennDiagram_1.6.20
## [101] regioneR_1.22.0 openssl_1.4.4
## [103] munsell_0.5.0 askpass_1.1
```